₹> d ibib abs hitstr 1-10

L5 ANSWER 1 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: DOCUMENT NUMBER:

2004:80450 CAPLUS

DOCOME

140:145835

TITLE:

Preparation of dibenzofused bicyclo[2.2.2]octanederived amides as modulators of the glucocorticoid

receptor

INVENTOR(S):

Vaccaro, Wayne; Yang, Bingwei Vera; Kim, Soong-hoon; Huynh, Tram; Tortolani, David R.; Leavitt, Kenneth J.;

Li, Wenying; Doweyko, Arthur M.; Chen, Xiao-tao;

Doweyko, Lidia

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Company, USA; et al.

SOURCE:

PCT Int. Appl., 265 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO. KI			KI	ND	DATE APPLICATION NO. DATE												
	WO 2004009017			A.	2 20040129			WO 2003-US22300 20030717										
		W:	ΑE,	AG,	ΑL,	AM,	AT,	ΑÚ,	ΑZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	ΊL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	OM,
			PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	ТĴ,	TM,	TN,
			TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW,	AM,	AZ,	BY,
					MD,												•	•
		RW:	GH,	GM,	ΚE,	ĻS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AT,	BE.	BG.
			CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC.
			ΝL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN.	GO.
			GW,	ML,	MR,	NE,	SN,	TD,	TG				·	·	•	•	•	~-
PRIORITY APPLN. INFO.:				. :				. 1	JS 20	002-3	3968	77:P	P :	2002	718			
OTHER GI	R SC	URCE	(S):			MAR	PAT 1	140:1	L4583	35							*	

II

AB Title compds. I [R-R4 = H, alk(en/yn)yl, alkoxy, aryl, etc.; Z = carboxamido, alkylamino, etc.] are prepared For instance, 2-amino-4,5-dimethylthiazole is coupled to the acid derived from the cycloaddn. of methacrylic acid and anthracene (CH3CN, EDCI, Et3N, HOAt, 18 h) to give II. I are glucocorticoid receptor modulators which are useful in treating diseases requiring glucocorticoid receptor agonist or antagonist therapy such as obesity, diabetes, inflammatory and immune disorders.

IT 312317-98-9P 650625-43-7P 650625-55-1P

Т

ANSWER 2 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN L5

ACCESSION NUMBER:

2004:80449 CAPLUS

DOCUMENT NUMBER:

140:157927

TITLE:

Homology modeling of nuclear hormone receptor Site II

and design of Site II ligands

INVENTOR(S):

Doweyko, Arthur; Nadler, Steven G.

PATENT ASSIGNEE(S):

Bristol-Myers Squibb Company, USA PCT Int. Appl., 276 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

650625-68-6P 650625-70-0P 650625-72-2P 650625-74-4P 650625-75-5P 650625-76-6P

DATE

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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PATENT NO.
                        KIND
                                               APPLICATION NO.
                                                                  DATE
     WO 2004009016
                         A2
                               20040129
                                               WO 2003-US22299
                                                                  20030717
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM,
              PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN,
              TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG,
              CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC,
              NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA,
              GW, ML, MR, NE, SN, TD, TG
PRIORITY APPLN. INFO.:
                                            US 2002-396907P X 20020718
     A binding site in nuclear hormone receptors is described and its
AΒ
     structural coordinates are provided. The invention provides
     machine-readable data storage media comprising structure coordinates of
     Site II and computer systems comprising the machine-readable data storage
             The invention provides methods used in the design and
     identification of ligands of Site II and of modulators of nuclear hormone
     receptors. The invention provides ligands of Site II, modulators of NHRs,
     pharmaceutical compns. comprising modulators of NHRs, methods of
     modulating NHRs, and methods of treating diseases by administering
     modulators of an NHR. Also provided are methods of designing mutants,
     mutant NHRs, Site II binding assays, and models of Site II.
ΙT
     312317-98-9P 650625-43-7P 650625-47-1P
     650625-55-1P 650625-58-4P 650625-60-8P
     650625-62-0P 650625-64-2P 650625-67-5P
```

Absolute stereochemistry.

5 ANSWER 3 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 2002:501478 CAPLUS

DOCUMENT NUMBER: 137:229148

TITLE: In vitro increase in chloroquine accumulation induced

by dihydroethano- and ethenoanthracene derivatives in

Plasmodium falciparum-parasitized erythrocytes

AUTHOR(S): Pradines, Bruno; Alibert, Sandrine; Houdoin, Carole;

Santelli-Rouvier, Christiane; Mosnier, Joel; Fusai, Thierry; Rogier, Christophe; Barbe, Jacques; Parzy,

Daniel

CORPORATE SOURCE: Unite de Parasitologie, Institut de Medecine Tropicale

du Service de Sante des Armees, Institut Federatif de

la Recherche 48, Marseille, 13998, Fr.

SOURCE: Antimicrobial Agents and Chemotherapy (2002), 46(7),

2061-2068

CODEN: AMACCQ; ISSN: 0066-4804

PUBLISHER: American Society for Microbiology

DOCUMENT TYPE: Journal LANGUAGE: English

had an amido group.

The effects of a series of dihydroethano- and ethenoanthracene derivs. on chloroquine (CQ) accumulation in CQ-susceptible strain 3D7 and CQ-resistant clone W2 were assessed. The levels of CQ accumulation increased little or none in CQ-susceptible strain 3D7 and generally increased markedly in CQ-resistant strain W2. At 10 μM , 28 compds. yielded cellular accumulation ratios (CARs) greater than that observed with CQ alone in W2. At 10 μ M, in strain W2, 21 of 31 compds. had CQ CARs two or more times higher than that of CQ alone, 15 of 31 compds. had CQ CARs three or more times higher than that of CQ alone, 13 of 31 compds. had CQ CARs four or more times higher than that of CQ alone, and 9 of 31 compds. had CQ CARs five or more times higher than that of CQ alone. At 1 $\mu\text{M}\text{,}~17~\text{of}~31~\text{compds.}$ had CQ CARs two or more times higher than that of CQ alone, 12 of 31 compds. had CQ CARs three or more times higher than that of CQ alone, 6 of 31 compds. had CQ CARs four or more times higher than that of CQ alone, and 3 of 31 compds. had CQ CARs five or more times higher than that of CQ alone. At 1 μM , 17 of 31 compds. were more potent inducers of CQ accumulation than verapamil and 12 of 31 compds. were more potent inducers of CQ accumulation than promethazine. The nature of the basic group seems to be associated with increases in the levels of CQ accumulation. At 1 and 10 μM , 10 of 14 and 13 of 14 compds. with amino group (amines and diamines), resp., had CARs ≥ 3 , while at 1

and 10 μ M, only 1 of the 13 derivs. with amido groups had CARs \geq 3. Among 12 of the 31 compds. which were more active inducers of CQ accumulation than promethazine at 1 μ M, 10 had amino groups and 1

448958-12-1, BG 1050 **448958-13-2**, BG 1051 **448958-14-3**, BG 1049

RL: BSU (Biological study, unclassified); BIOL (Biological study) (in vitro increase in chloroquine accumulation induced by dihydroethano- and ethenoanthracene derivs. in Plasmodium falciparum-parasitized erythrocytes)

RN 448958-12-1 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-2-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 448958-13-2 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-3-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 448958-14-3 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-4-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT:

THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 4 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN ACCESSION NUMBER: 2002:462893 CAPLUS

10/621,909

DOCUMENT NUMBER:

137:185297

TITLE:

Synthesis and Effects on Chloroquine Susceptibility in

Plasmodium falciparum of a Series of New

Dihydroanthracene Derivatives

AUTHOR(S):

Alibert, Sandrine; Santelli-Rouvier, Christiane; Pradines, Bruno; Houdoin, Carole; Parzy, Daniel; Karolak-Wojciechowska, Janina; Barbe, Jacques

CORPORATE SOURCE:

GERCTOP-UMR CNRS 6009, Faculte de Pharmacie,

Universite de la Mediterranee, Marseille, 13385, Fr. Journal of Medicinal Chemistry (2002), 45(15),

SOURCE: 3195-3209

CODEN: JMCMAR; ISSN: 0022-2623

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 137:185297

To suggest a mechanism of action for drugs capable of reversing the AΒ chloroquine resistance in Plasmodium falciparum, a new set of 9,10-dihydro-9,10-ethano- and -ethenoanthracene derivs. was synthesized and tested to assess their effect on chloroquine susceptibility in resistant strains of Plasmodium falciparum. With respect to this, reversal of resistance and change in drug accumulation were compared. Structure-activity relationship and mol. modeling studies made it possible to define a pharmacophore moiety for reversal agents and to propose a putative model of interaction with some selected amino acids.

ΊΤ 448958-12-1P 448958-13-2P 448958-14-3P

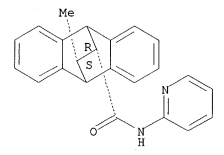
> RL: PAC (Pharmacological activity); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(preparation and effects on chloroquine susceptibility in Plasmodium falciparum of a series of ethano- and ethenodihydroanthracenes)

448958-12-1 CAPLUS RN

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-2-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.



RN 448958-13-2 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-3-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 448958-14-3 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-12-methyl-N-4-pyridinyl-, (11R,12S)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT:

48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 5 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1999:220218 CAPLUS

DOCUMENT NUMBER:

130:237374

TITLE:

New tetracyclo[6.6.2.02,7.09,14]hexadeca-

2(7), 3, 5, 9(14), 10, 12] hexaenes as phospholipase

inhibitors

INVENTOR(S):

Friebe, Walter-Gunar; Tibes, Ulrich; Scheuer, Werner

PATENT ASSIGNEE(S): Roche Diagnostics G.m.b.H., Germany

SOURCE:

Ger. Offen., 10 pp.

DOCUMENT TYPE:

CODEN: GWXXBX

DOCOMENT I

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APF	LICATION NO.	DATE				
DE 19742014	Al 19990	0325 DE	1997-19742014	19970924				
ZA 9808711	A 20000	0323 ZA	1998-8711	19980923				
CA 2304879	AA 19990	0401 CA	1998-2304879	19980924				
WO 9915493	A1 19990	0401 WO	1998-EP6096	19980924				
W: AL, AM,	AT, AU, AZ,	BA, BB, BG, B	BR, BY, CA, CH,	CN, CU, CZ, DE,				
				IS, JP, KE, KG,				
KP, KR,	KZ, LC, LK,	LR, LS, LT, L	U, LV, MD, MG,	MK, MN, MW, MX,				
NO, NZ,	PL, PT, RO,	RU, SD, SE, S	G, SI, SK, SL,	TJ, TM, TR, TT,				
UA, UG,	US, UZ, VN,	YU, ZW, AM, A	Z, BY, KG, KZ,	MD, RU, TJ, TM				
				CY, DE, DK, ES,				
FI, FR,	GB, GR, IE,	IT, LU, MC, N	L, PT, SE, BF,	BJ, CF, CG, CI,				
CM, GA,	GN, GW, ML,	MR, NE, SN, T	D, TG					

AU 9897466 A1 AU 1998-97466 19980924 19990412 20000829 BR 1998-13217 19980924 BR 9813217 Α EP 1998-951464 19980924 20000913 EP 1034162 A1 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE, FI Т2 TR 200001222 20000921 TR 2000-20000122219980924 JP 2002505999 T2 20020226 JP 2000-512804 19980924 PRIORITY APPLN. INFO .: DE 1997-19742014 A 19970924 WO 1998-EP6096 W .19980924

OTHER SOURCE(S):

MARPAT 130:237374

GI

$$X$$
 ZY R^2 I

AB Title compds. I [R1, R2 = H, halogen; X = H, Y = (un)substituted NH2, N+H2Me; XY = (un)substituted CH2NH; Z = CH2, C:NH] were prepared for use as phospholipase inhibitors (no data). Thus, I [R1, R2, X = H, Y = NH2.HCl, Z = CH2] was treated with KOCN to give I [R1, R2, X = H, Y = NHCONH2, Z = CH2].

IT 221352-72-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of tetracyclohexadecahexaenes as phospholipase inhibitors)

RN 221352-72-3 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 9,10-dihydro-N-4-pyridinyl- (9CI) (CA INDEX NAME)

IT 221352-19-8P 221352-21-2P 221352-47-2P

221352-50-7P 221352-54-1P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of tetracyclohexadecahexaenes as phospholipase inhibitors)

RN 221352-19-8 CAPLUS

CN 4-Pyridinamine, N-[(9,10-dihydro-9,10-ethanoanthracen-11-yl)methyl]- (9CI) (CA INDEX NAME)

10/621,909

RN 221352-21-2 CAPLUS

CN 4-Pyridinamine, N-[(1,5-dichloro-9,10-dihydro-9,10-ethanoanthracen-11-yl)methyl]- (9CI) (CA INDEX NAME)

RN 221352-47-2 CAPLUS

CN 1H-Imidazol-2-amine, N-[(1,5-dichloro-9,10-dihydro-9,10-ethanoanthracen-11-yl)methyl]-4,5-dihydro- (9CI) (CA INDEX NAME)

RN 221352-50-7 CAPLUS

CN 1H-Imidazol-2-amine, N-[(9,10-dihydro-9,10-ethanoanthracen-11-yl)methyl]-4,5-dihydro-(9CI) (CA INDEX NAME)

$$CH_2-NH$$

RN 221352-54-1 CAPLUS

CN 1H-Imidazol-2-amine, N-[(4,5-dichloro-9,10-dihydro-9,10-ethanoanthracen-11-yl)methyl]-4,5-dihydro-, monohydrobromide (9CI) (CA INDEX NAME)

HBr

L5 ANSWER 6 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1995:457943 CAPLUS

DOCUMENT NUMBER: 123:199069

TITLE: Infrared spectra of N-ferrocenylamic acids

AUTHOR(S): Yang, Bingqin; Ma Huairang; Chen Zhibin

CORPORATE SOURCE: Department of Chemistry, Northwest University, Xian,

710069, Peop. Rep. China

SOURCE: Guangpuxue Yu Guangpu Fenxi (1995), 15(1), 53-6

CODEN: GYGFED; ISSN: 1000-0593

PUBLISHER: Beijing Daxue Chubanshe

DOCUMENT TYPE: Journal

LANGUAGE: Chinese

AB The IR spectra of seven N-ferrocenylamic acids are reported. The effect

of intramol. hydrogen bonds was discussed and the absorption characteristics of some structural isomers were reported. In addition, the

mol. structures of some interconversion isomers were proven by IR.

IT 146083-59-2

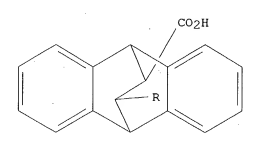
RL: PRP (Properties)

(IR spectra of N-ferrocenylamic acids)

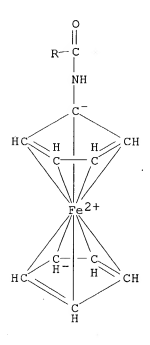
RN 146083-59-2 CAPLUS

CN Ferrocene, [[(12-carboxy-9,10-dihydro-9,10-ethanoanthracen-11-yl)carbonyl]amino]-, cis- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A



L5 ANSWER 7 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1994:244394 CAPLUS

DOCUMENT NUMBER:

120:244394

TITLE:

Dibenzo-fused derivatives of bicyclo[2.2.2]octane as

cholecystokinin inhibitors

INVENTOR(S):

Kalindjian, Sarkis Barret; Low, Caroline Minli Rachel;

Mcdonald, Iain Mair; Hull, Robert Antony David; Shankley, Nigel Paul; Buck, Ildiko Maria; Steel, Katherine Isobel Mary; Davies, Jonathan Michael

Richar; Dunstone, David John; et al.

PATENT ASSIGNEE(S):

SOURCE:

James Black Foundation Ltd., UK

PCT Int. Appl., 159 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	CENT I	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	ο.	DATE				
WO	9316982		A1 199		19930902			WO 1993-GB346			19930219							
	W:	AT,	AU,	BB,	BG,	BR,	CA,	CH,	CZ,	DE,	DK,	ES,	FI,	GB,	HU,	JP,	KP,	
		KR,	ΚZ,	LK,	LU,	MG,	MN,	MW,	NL,	NO,	NZ,	PL,	PΤ,	RO,	RU,	SD,	SE,	
		SK,	UA,	US														
	RW:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	ΙE,	ΙΤ,	LU,	MC,	NL,	PT,	SE,	
		BF,	ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	ML,	MR,	SN,	TD,	TG				
ΑU	9335	097		A.	1	1993	0913		Α	J 19	93-3	5097		1993	0219			
	9301											193						
	6269					1994	1207		E	P 19	93-9	04230)	1993	0219			
EΡ	6269	42		B	1	1997	0423											
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JΡ	0750											14633				·	•	
HU	71499	9		Αź	2	1995	1128		н	J 19	94-2	280		1993	0219			
AT	15209	95		E		1997	0515		A.	Ր 19:	93-9	04230)	1993	0219			

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US 5514683
                             19960507
                                                               19940809
                        Α
                                             US 1994-288185
     NO 9403055
                        Α
                             19941011
                                             NO 1994-3055
                                                               19940818
     FI 9403817
                             19940819
                                             FI 1994-3817
                                                               19940819
                        Α
PRIORITY APPLN. INFO.:
                                          GB 1992-3608
                                                               19920220
                                          GB 1992-13093
                                                            Α
                                                               19920619
                                          GB 1992-24629
                                                            A
                                                               19921124
                                          WO 1993-GB346
                                                            Α
                                                               19930219
                                          GB 1993-16722
                                                               19930812
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OTHER SOURCE(S):

MARPAT 120:244394

Ι

GΙ

$$\begin{array}{c|c} & & & & \\ & & & & \\ ZW & & & & \\ R1 & & & \\ R6 & & \\ R6 & & \\ R6 & & & \\ R6 & & \\ R6$$

Title compds. I [W = CO, SO, SO2; X = CO, SO, SO2, COCH2 (with CO end AB bound to Y), provided that ≥1 of W and X contains CO; Y = certain (un) substituted OH or NH2 groups; Z = different (un) substituted OH or NH2 groups; R1 = H, Me, halo, (amidated or esterified) CO2H or CH2CO2H; R2 = groups for R1, or COZ' (Z' = Z) when Z is absent and W = H; or R1R2 = pi bond; R3, R4 = halo, amino, NO2, cyano, SO2NH2, alkyl, alkoxy, (amidated or esterified) CO2H; R5, R6 = H, R3; m, n = 0-4, provided that both are \leq 2.unless R3 or R4, resp., are exclusively halo] were prepared as ligands binding at cholecystokinin (CCK) and gastrin receptors. Thus, 2,3,5,6-dibenzobicyclo[2.2.2]octane-7,8-dicarboxylic acid anhydride reacted with 1-adamantanemethylamine, the resultant acid-amide was condensed with H-L-Pro-OCH2Ph.HCl using PyBOP, and the benzyl ester function was hydrogenolyzed and reesterified with diazomethane to give title compound cis-II as a mixture of 2 diastereomers which were separated by repeated crystallization These isomers bound to CCKB receptors (mouse cortical membrane) with pKi = 5.8 and 7.3. Included are 238 synthetic examples, 1H NMR data for all final products (free bases or N-methyl-D-glucamine salts), and receptor-binding results (CCKA, CCKB, and gastrin) for most I. IT 153459-03-1P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of, as CCK and gastrin antagonist)

RN 153459-03-1 CAPLUS

2-Furancarboxylic acid, 5-[[[9,10-dihydro-12-[[(tricyclo[3.3.1.13,7]dec-1-ylmethyl)amino]carbonyl]-9,10-ethanoanthracen-11-yl]carbonyl]amino]-,

methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

ANSWER 8 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1993:102173 CAPLUS

DOCUMENT NUMBER:

118:102173

TITLE:

Synthesis and reaction of N-ferrocenylamic acid

AUTHOR(S):

Ma, Huairang; Chen, Zhibing; Yang, Bingqin; Li,

CORPORATE SOURCE:

Dep. Chem., Northwest Univ., Xian, 710069, Peop. Rep.

China

SOURCE:

Gaodeng Xuexiao Huaxue Xuebao (1992), 13(5), 633-5

CODEN: KTHPDM; ISSN: 0251-0790

DOCUMENT TYPE:

LANGUAGE:

Journal

Chinese

GΙ

AΒ Seven N-ferrocenylamic acids have been synthesized and characterized by elemental anal., IR and 1H NMR. The investigation on the chelation of acid I (Fc = ferrocenyl) with Cu2+ and Zn2+ ions demonstrates the coordination of the oxygen atom on the amide group to the metallic ion.

IT 146083-59-2P

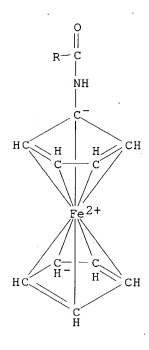
> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 146083-59-2 CAPLUS

CN Ferrocene, [[(12-carboxy-9,10-dihydro-9,10-ethanoanthracen-11yl)carbonyl]amino]-, cis- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A



ANSWER 9 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

DOCUMENT NUMBER:

1989:149699 CAPLUS 110:149699

TITLE:

Solid precursors, methods, and apparatus for

development of latent fingerprints with cyanoacrylates

INVENTOR(S):

Warrener, Ronald Norman; Yong, Siaw Jan

PATENT ASSIGNEE(S):

Australian National University, Australia

SOURCE:

PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 8801616	A1	19880310	WO 1987-AU286	19870825

W: AU, JP, US

RW: AT, BE, CH, DE, FR, GB, IT, LU, NL, SE

AU 8779188 A1 19880324 AU 1987-79188 19870825 PRIORITY APPLN. INFO.: AU 1986-7669 19860825 WO 1987-AU286 19870825

AB Solid precursors of cyanoacrylate monomers, such as Diels-Alder adducts, homopolymers, or cyclic or acyclic precursors, may be used in a method for the development of a latent fingerprint on a surface by exposure of the surface to cyanoacrylate monomer generated from a solid precursor to form a cyanoacrylate-developed print. Preferably, the solid precursor includes colored or fluorescent substituent groups. An apparatus for use in the cyanoacrylate development of latent fingerprints is diclosed also. A mixture of anthracene, Super Glue, hydroquinone, and dry benzene was refluxed at 90° for 12 h. The anthracene/Et 2-cyanoacrylate adduct was treated with NaOH to make anthracene/2-cyanoacrylic acid adduct (m. 206-209°) in 99.7% yield.

IT 119858-97-8

RL: PRP (Properties)
(NMR of)

RN 119858-97-8 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 11-cyano-9,10-dihydro-N-8-quinolinyl-(9CI) (CA INDEX NAME)

IT 119859-04-0 119859-05-1

RL: BIOL (Biological study)

(as solid precursor for cyanoacrylate development of latent

fingerprints)
RN 119859-04-0 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxamide, 11-cyano-9,10-dihydro-N-(4-methyl-2-oxo-2H-1-benzopyran-7-yl)- (9CI) (CA INDEX NAME)

CN 9,10-Ethanoanthracene-11-carboxamide, 11-cyano-9,10-dihydro-N-(2-oxo-2H-1-benzopyran-7-yl)- (9CI) (CA INDEX NAME)

L5 ANSWER 10 OF 10 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER: 1979:120619 CAPLUS

DOCUMENT NUMBER: 90:120619

TITLE: Reactions of aminopyridines with some inner anhydrides

AUTHOR(S): El-Zanfally, S.; El-Basil, S.

CORPORATE SOURCE: Org. Chem. Dep., Fac. Pharm., Cairo, Egypt

SOURCE: Egyptian Journal of Pharmaceutical Sciences (1978),

Volume Date 1976, 17(1), 53-62

CODEN: EJPSBZ; ISSN: 0301-5068

DOCUMENT TYPE: Journal LANGUAGE: English

AB Reaction of 3-aminopyridine inner anhydrides, e.g., succinic or maleic anhydride, gives amic acids. In the case of 2- and 4-aminopyridines, the products obtained were amic acids or cyclized products (succinimide derivs.). With maleic anhydride, both 2- and 4-aminopyridines form a charge-transfer complex. Phenylsuccinic anhydride behaves as a C acid with 4- but not with 2-aminopyridine.

IT 69537-47-9P 69537-51-5P

RN 69537-47-9 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxylic acid, 9,10-dihydro-12-[(2-pyridinylamino)carbonyl]-, cis- (9CI) (CA INDEX NAME)

RN 69537-51-5 CAPLUS

CN 9,10-Ethanoanthracene-11-carboxylic acid, 9,10-dihydro-12-[(4-pyridinylamino)carbonyl]-, cis- (9CI) (CA INDEX NAME)

10/621,909

=> d his

(FILE 'HOME' ENTERED AT 14:10:23 ON 19 MAY 2004)

FILE 'REGISTRY' ENTERED AT 14:10:31 ON 19 MAY 2004

L1 STRUCTURE UPLOADED

L2 25 S L1

L3 588 S L1 FULL

FILE 'CAPLUS' ENTERED AT 14:11:24 ON 19 MAY 2004

L4 2 S L3/THU

L5 10 S L3

=> d 11

L1 HAS NO ANSWERS

L1 STR

Structure attributes must be viewed using STN Express query preparation.

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